Remarks

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This Amendment responds to the Office Action dated February 27, 2003. In the Office Action, the examiner objected to claims 7 and 8 due to an informality regarding recitation of a Markush group. Claims 7 and 8 have been amended to refer to "the group consisting of" as requested by the patent examiner.

In the Office Action, the examiner rejected claim 6 under 35 U.S.C. §112, second paragraph as being indefinite, particularly relating to the word "composite." Claim 6 has been amended to delete that word such that the claim now provides "dyeing the laminated fabric of said knit layer, aluminum sheet layer and non-woven fabric layer with a background color." Accordingly, applicant has overcome the indefinite use of "composite" and respectfully requests that the examiner withdraw this rejection of claim 6.

In the Office Action, the examiner rejected as being non-patentable claim 1 under 35 U.S.C. §102(b) in view of U.S. Patent No. 5,149,582 to LaMarca (herein "LaMarca '582"). The examiner rejected claims 2-5 and 10-11 under 35 U.S.C. §103(a) in view of U.S. Patent No. 4,901,738 to Brink (herein "Brink '738"). The examiner rejected claim 6 under Section 103(a) as being unpatentable over LaMarca '582; Brink '738 and U.S. Patent No. 5,568,666 to Seibert. The examiner rejected claims 7-9 and 12 under 35 U.S.C. §103(a) as being unpatentable in view of LaMarca '582; Brink '738 and U.S. Patent No. 3,094,547 to Heine ("Heine '547").

Further, the examiner rejected under the doctrine of obviousness-type double patenting claims 1-12 as being unpatentable over claims 1-12 and 19-21 of co-pending application by applicant Principe, Serial No. 09/872,942. Applicant respectfully submits that she will file a Terminal

Disclaimer thereby avoiding the obviousness-type double patenting rejection upon allowance of claims in both of these patents applications.

Applicant has amended claim 1 to recite "A cigar and cigarette burn resistant gaming table cloth which resists decomposition by burn and char formation by a lit cigarette or cigar for at least three minutes during direct contact exposure to a the lit end of the eigar or eigarette." Further, claim l has been amended to provide "said aluminum acting as a heat sink and preventing decomposition by burn and char formation on the fabric for at least three minutes."

Support for these concepts is found throughout the patent specification. Particularly in paragraph 3, "Casino patrons dislike the look of stained gaming table covers (originating from spilled drinks) and table covers with burn marks are similarly unsightly. Patrons will avoid these stained and abused gaming tables ... the distressed look of the table engenders a poor quality image in the minds of casino patrons." "Hence, there is a need for a water resistant cigar and cigarette burn resistant gaming table of billiard table cover." Paragraph 5.

In other words, applicant seeks protection for a cigarette burn resistant cloth that prevents decomposition by burn and prevents char formation due to direct contact exposure to a lit cigarette for at least three minutes.

It is respectfully submitted that the references cited by the examiner are primarily concerned with flame resistant or flame retardant fabrics whereas the presently claimed invention is more concerned with a burn resistant fabric which has been more specifically identified as "preventing decomposition by burn and char formation on the fabric for at least three minutes." In Paragraph 13, it is an object of the invention to provide "a cigar and cigarette burn resistant gaming table cloth." In Paragraph 15, the gaming table cloth "significantly resists burning by a lit cigarette or a lit cigar cigarette." Paragraph 22 discusses experiments indicating that prior art gaming table covers burn in 11-20 seconds upon direct contact with lit cigarette or lit cigar which have an approximate temperature of 1200°F. "The cloth composite of the present invention does not burn unless the hot spot exposure exceeds at least 3 minutes of direct contact with a lit cigarette or lit cigar. It is believed that the aluminum acts as a heat sink and prevents burn on the fabric for up to 3-4 minutes."

In view of these explicit statements in the patent specification, the present invention is not primarily concerned with flame retardant fabrics but rather is concerned with burn resistant fabrics wherein burn resistant is further defined as "preventing decomposition by burn and char formation on the fabric for at least three minutes." The entire purpose of the new fabric was to avoid burn marks due to cigars and cigarettes on gaming and billiard table cloth. See Paragraph 3 (casino patrons dislike ... table covers with burn marks); Paragraph 5 (there is a need for water resistant cigar and cigarette burn resistant gaming table); Paragraph 13 (same); Paragraph 15 (a cloth which significantly resists burning by a lit cigarette or a lit cigar for about three minutes); and Paragraph 22 (the present cloth invention does not burn unless the hot spot exposure from the lit cigarette or lit cigar exceeds at least three minutes of direct contact). Hence, the present invention is not directed to a flame retardant or flame resistant fabric but rather is directed to a fabric which prevents decomposition by burn and char formation (such burn or char formation representing burn marks on casino table cloth and billiard cloth per Paragraph 3).

Persons with ordinary skill in the art of burn resistant and flame retardant fabrics recognize the relationship between flame retardant features and char formation. Exhibit A attached provides a glossary for the Flame Retardants Center which glossary separately identifies char (carbonaceous

residue resulting from pyrolysis or incomplete combustion), scparate from fire retardant and fire retardants. (pp. A-2, A-3). Further the book Fire Retardency of Polymeric Materials, Ed. Arthur F. Grand and Charles A. Wilky, Pub. Marcel Dekker, Inc. (2000) pp. 172-174, (Exhibit B attached) discusses the role of char in the study of thermal degradation in fabrics. p. B-3. In Exhibit C attached, persons of ordinary skills in the art recognize that the phrase "fire retardant" encompasses many factors including ease of ignition, flame spread, fire endurance, rate of heat release, case of extinction, smoke evolution and toxic gas evolution. Exhibit D enclosed also discusses char formation and the relationship of char formation regarding flame retardant polymers. Exhibit E attached is a book Flammability of Fabrics. Charles Hilado, Technomic Pub. Co., Westport, CT (1974) pp. 11-24 and 59, 62 and 63, which discusses char life at p. E-4 and the production of char at p. E-5 and states that many flame retardants have a tendency to produce more char at pp. E-7 and E-8. The article "Flammability Measurements and Thermal Decomposition of Textiles" by J.E. Hendrix, et. al. and that same textbook Flammability of Fabrics, at p. E-11 graphically illustrates at p. E-12 that fabrics that produce more char have lower combustion temperatures as compared with fabrics that are not treated with flame retardant chemicals.

In other words, (a) production of char is a recognized characteristic in the flame retardant industry, that is, recognized by the person of ordinary skill in the art, and (b) typically char formation would be increased in flame retardant fabrics rather than prevented or reduced as claimed in the present invention. Further, the present invention is interested in preventing both burn and char formation for at least three minutes after direct contact exposure to a lit end of a cigar or cigarette. These attached references, and the fire retardant prior art references cited by the examiner, teach

away from the claimed invention which prevents decomposition by burn and prevents char formation on fabric for three minutes due to the direct contact exposure to the lit end of a eigarctte.

None of the references cited by the examiner discuss <u>preventing decomposition</u> by burn and char formation on the fabric due to direct contact with a lit eigarette or eigar for at least three minutes as presently claimed in the present invention. The patent specification, and now the amended claims, are all directed to preventing decomposition by burn and decomposition by char formation. The articles attached hereto indicate that char formation is one of many physical characteristics which have an impact on the flame retardant feature of fabrics. However, the technology, at least as explained in Exhibits D and E indicates that fabrics which generate greater char have greater flame retardant characteristics. In the present invention, char formation and burn decomposition is avoided (because casino patrons dislike table covers with burn marks, Paragraph 3). Any resulting flame retardant characteristic of the gaming table cloth is, in the present patent application, irrelevant to the structure and the operation of the structure upon "direct contact exposure to a lit end of the eigar or eigarette."

LaMarca '582 is specifically directed to a flame barrier, puncture resistant fabric sheet as stated in its title. Flammability requirements are identified at column 1, line 25-30 and lines 35-49. LaMarca '582 specifically states that these flame tests involve a horizontal type flame test and a vertical type burn test and that "flame must be prevented from penetrating through the upholstery fabric." Col. 1, line 36. LaMarca '582 discusses a flame barrier composite sheet material at col. 1, line 66 and col. 2, line 3. In discussing the aluminum foil layer at col. 4, line 36, LaMarca '582 states "The metal foil provides a flame barrier and also serves as a heat sink in the resulting composite fabric cheap material and aids in stopping vertical flame propagation." Lines 40-43.

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Further, at col. 5, lines 24-30, it is explained "It is recognized that the composite fabric sheet material of the [LaMarca '582] invention will have applications in a wide variety of fields where a highly flame and puncture resistant fabric sheet material is desired." In explaining the preferred embodiment of his invention, LaMarca '582 discusses the flammability specifications and fire block test at col. 7, lines 4-6. LaMarca '582 does not discuss preventing decomposition by burn nor decomposition by char formation.

With respect to Brink '738, that patent disclosure discusses a laser shield which claims are particularly directed to a laser resistant surgical drape. The laser shield claim format in Brink '738 is a similar claim format in that the present claim relates to a gaming table cloth which resists decomposition by burn and char formation by a lit cigarette or cigar for at least three minutes. The structural aspect of preventing both (i) decomposition by burn and (ii) preventing char formation for at least three minutes is explicitly recited in claim 1 of the present invention. In other words, the flame resistant fabric identified in LaMarca '582 is directed to a different composite cloth structure as compared with the presently claimed invention which is directed to a gaming table cloth which resists unsightly burn marks caused by a lit cigar or cigarette. Brink '738 discusses a need for a laser resistant surgical drape at col. 1, line 16 and line 59, col. 2, lines 16 and 48. More particularly, the laser shield in Brink 738 includes thermal plastic fibers which resist flaming when subjected to a carbon dioxide laser. Col. 4, lines 17-22. More significantly the laser shield in Brink '738 uses aluminum or many other metal foils such as copper, gold, iron, and steel. Col. 4, lines 40-50. In the present invention, other metal foils or sheets have been found to be unacceptable for the burn resistant fabric. Paragraph 48(steel does not adhere to cloth layers and is too heavy).

Brink '738 also discusses a national fire standard which relates to ignition in 15 seconds or 20 seconds. Col. 7, lines 17-25. In direct contrast, the presently claimed invention resists decomposition by burn and resists decomposition by char formation for at least three minutes due to direct contact exposure to the lit end of a cigar or cigarette. The claimed invention describes a different structure for a different purpose which, most importantly, operates in a different manner when subjected to "direct contact exposure." The claimed prevention of char (and the resulting burn marks, Paragraphs 3, 13, 15 and 22) is not disclosed by the prior art. In fact, the prior art Exhibits D and E attached show that greater char formation enhances fire retardant features of a fabric.

Scibert '666 and Heine '547 do not show, teach or suggest a laminated fabric with an aluminum sheet which prevents decomposition by burn and char formation on a fabric for at least three minutes during direct contact exposure to lit end of a cigar or cigarette.

Accordingly, it is respectfully requested that the examiner withdraw the rejection applied against claims 1-12 and allow those claims in the present case.

Respectfully submitted,

Fleit, Kain, Gibbons, Gutman & Bongini, P.L.

June 17,2003

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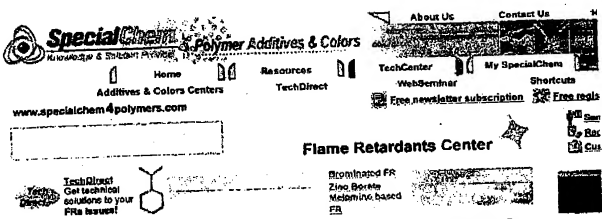
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Glossary

DSBG

Testino

Why use Flame Retardents ?

Families of Flame Reterdants

Regulation and Testing Fire Dynamics Building & Construction Electrical Engineering Transportation Furniture

Future Trends **▶** Glossary Tochnical Library This glossary is partially inspired from the ISO 13943: 2000 -fire safety vocabulary-

A-E F G-N O-T

Select a

after flame

A flame which persists after the ignition source has been removed.

after flame time (s)

Length of time for which an after flame persists under specified conditions.

Persistence of glowing efter cessation of flaming or if no flaming occurs, after the ignition source have been removed

afterglow time (\$)

The length of time for which an afterglow persists under specified test conditions.

The ability of a material to resist the influence of an electric arc along its surface, under specified test conditions

burned area (m²)

That part of the damaged area of a material that has been destroyed by combustion or pyrolysis, under specified conditions. (See also damaged area - N 27)

burned length (m)

The maximum length of a material that has been destroyed by combustion or pyrolysis under specified conditions.

burning behavlour

All the physical and/or chemical changes that take place when an item is exposed to a specified ignition source.

calorific value (J/kg)

Heat produced by combustion of unit mass of a given substance.

gross calorific value (J/kg)

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Exhibit A

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The calorific value of a substance when the combustion is complete and any produced water is entirely condensed under specified conditions.

net calorific value; heat of combustion (J/kg) The calorific value of a substance when the combustion is complete and any produced water is in the vapour state under specified conditions.

char (noun)

Carbonaceous residue resulting from pyrolysis or incomplete combustion.

combustible (adj.)

Capable of being combustible.

combustion

Exothermic reaction of a combustible substance with an oxidizer.

nodergation

A combustion wave, accompanied by an explosion, propagating at subsonic velocity.

A combustion wave, accompanied by an explosion, propagating at supersonic velocity and characterized by a shock wave.

end use conditions

The intended conditions to which an item will be subjected during its normal working life, when used in accordance with the manufacturer's instructions.

explosion

An abrupt expansion of gas which may result from a rapid exidation or decomposition reaction, with or without an increase in temperature.

exposure time (s)

The length of time for which people, animals or items are exposed under specified test conditions.

A-E E G-N O-I

Self supporting combustion spreading uncontrolled in time and space characterized by the emission of heat and effluent often accompanied by flame and/or glowing.

fire barrier (function).

The ability of a separating element to resist the passage of flame and/or heat-and/or effluents.

Fire.barrier (product)

A separating element able to function as a fire barrier (function), for a period of time under specified conditions.

fire behaviour

The change in the physical and/or chemical properties of an item and/or structure exposed fire,

fire compartment

An enclosed space, which may be subdivided, bounded by elements of

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construction having a specified fire resistance.

meuffle enn

The totality of gases, and/or aerosols (including suspended particles) created by combustion or pyrolysis.

fire exposure

The extent to which persons, animals or items are subjected to the conditions created by fire.

fire gases

The gaseous part of the products of combustion.

The potential for injury and/or damage from fire.

The response of an item when exposed to a specific fire.

The ability of an item to fulfil for a stated period of time the required stability and/or integrity and/or thermal insulation and/or other expected duty specified in a standard fire resistance test.

fire retardant (noun)

A substance added or a treatment applied to a material in order to delay ignition or to reduce the rate of combustion:

fire acenario

A detailed description of conditions, including environmental, of one or more stages from before ignition to after completion of combustion in an actual fire at a specific location or in a real scale simulation.

fire simulation

Mathematical representation of one or more of different interconnected phenomena governing a fire development.

Zone of combustion in the gaseous phase usually with emission of light.

flame front

The boundary of flaming combustion at the surface of a material or propagating through a gaseous mixtura.

flame retardance

The property of a material whereby flaming combustions is solved terminated or prevented.

flame retardant (noun)

A substance added, or a treatment applied to a material in order to suppress or delay the appearance of a flame and/or reduce its propagation (spread) rate.

flame retardant treatment

A process whereby improved flame retardance is imparted to a material or product.

flame spread

Propagation of a flame front.

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flame spread rate (m/s). Distance travelled by a flame front during its propagation, divided by the time of travel, under specified conditions.

flammability
The shrifty of a material or product to burn with a flame under specified conditions.

nammable Capable of flaming under specified conditions.

mash-over The transition to a state of total surface involvement in a fire of combustible materials within an enclosure.

fully developed fire The spin-of this involvement of combustible ma

A-E E G-N Q-T

gasification Transformation of a solid and/or liquid material to a geseous state.

glowing.combustion Combustion of a material in the solid phase without flame but with emission of tight from the combustion zone.

gross calorific value (J/kg)
The calorific value of a substance when the combustion is complete and any produced water is entirely condensed under specified conditions.

heat flux (kW/m²) The amount of thermal energy emitted, transmitted or received per unit he amount of the

heat release (J). The calonitic energy which is released by the combustion of an item under specified conditions.

host release rate : rate of heat release (W/m2 or W) The calcrific energy released per unit lime by an item during combustion under specified comittions.

heat stress Conditions caused by exposure to elevated/reduced temperature, radiant rest flox, or combination of these factors.

ignitability filegass of the case with which an item can be ignited, under specified conditions.

Ignition Initiation of combustion,

ignition source A source of energy that initiates combustion.

incandescence Emission of light produced by a material when integely heated. It can be produced by materials in the light produced by the prod

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integrity
The integrity of a separating element when exposed to fire on one side, to
The integrity of a separating element when exposed to fire on one side, to
prevent the passage of firmes and hot gases on the occurrence of firmes
on the unexposed side, for a stellad period of time in a standard fire
resistance test...

intermediate scale test Test performed on an item of medium dimensions.

*L.C. 50 : Teth'al concentration 50, form3)
The concentration of toxic gas or fire effluent statistically calculated from concentration-response data to produce lethality in 50 % of test animals of given species, under specified conditions.

large scale test.
Test, which cannot be carried out in typical taboratory chamber, partierned on an item of large dimensions.

limit oxygen index (LOI) (%)
The minimum concentration of congen in a minime of congen and nitrogen that will just support fisming combustion of a material under specified conditions.

melting behaviour
The phenomena accompanying the softening of a material under the influence of heat (including deforming and dripping, but not fleming).

motten drips (noun): Falling droplats of material which have been softened or liquefied by

net calorific value : heat of combustion (J/kg)
The calorific value of a substance when the combustion is complete end
eny produced water is in the vapour state under specified conditions:

non-combustible

Not expecte of undergoing combustion under specified conditions.

non-flammable Not capable of burning with a flame under specified conditions.

ARE GEN OLI

opacity of smoke (dimensionless)
A measure of the attenuation of a light beam passing through emoke appressed as the ratio (FT) of incident luminous flux (T) to transmitted luminous flux (T) through smoke under specified conditions.

optical density of arroles

A measure of the attenuation of a light beam passing through smoke appressed as the common logarithm (i.e. logarithm to the base 10) (lg(VI)) of the opacity of smoke.

pitol(ed) ignition

Ignition of combustible gases or vepours by a secondary source of
Ignition of combustible gases or vepours by a secondary source of
energy, has by for exempte, a theme, speak, electrical and or glowing wire.

products of combustion The solid. Nguid and goscous materials resulting from combustions.

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That part of the ineversible chemical decomposition caused solely by a rise in temperature.

The response of a material in contributing by its own decomposition to a fire to which it is exposed, under specified conditions.

Test which simulates a given application, taking into account the real scale, the real way of working or installation and the environment.

A preliminary test used for ascertaining whether an item is likely to exhibit (or not) certain characteristics according to a standardized test method.

self-extinguishing Deprecated term.

small scale test

A test performed on an item of small dimensions.

amoke

Visible part of fire effluent.

emoke rescuration

The reduction, (usually expressed as a percentage), in the intensity of light due to its passage through smoke.

The combustion of a material without flame and without light being visible.

Particulate matter produced and deposited during or after combustion.

specific optical density of amoke (dimensionless)

A measure of the opacity of the smoke produced by a specimen taking irsts account the optical density and factors characteristic of the specified test method.

surface spread of flame.

The propagation of flame away from the source of ignition across the surface of a liquid or a solid:

sustained flaming

Persistence of flame on or over a surface for a minimum period of time.

thermal degradation

A process whereby the action of heat or elevated temperature on an item causes a loss of some properties e.g physical, mechanical or electrical.

The potential for injury or loss of life by exposure to toxicants with respect to their potency, 'quantity, concentration and duration of exposure,

A measure of the emount of toxicant required to effect a specific toxic effecE

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Transmittance (through smoke) The ratio of transmitted luminous flux (T) through smoke to incident turninous flux (1) under specified conditions.

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Fire Retardancy of Polymeric Materials

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NEW YORK · BASEL

Exhibit B B-1

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Levchik and Wilkle

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THE ROLE OF CHAR IN THERMAL DEGRADATION: FIRE RETARDANCY

Char, which is formed in the process of thermal degradation, can play several roles in fire retardancy. The formation of char in and of itself has a significant effect on the degradation because that fortuation must occur at the expense of other reactions that may form volatiles; thus, char formation may limit the amount of fuci available. An example of this occurs in cellulose, which may degrade either by a series of dehydration reactions that yield water, carbon dioxide, and char. or by a process in which levoglucosan is produced, which eventually leads to the formation of volatiles (1); this is shown as Scheme 1.

Scheme 1 Thermal degradation of cellulose to char or flammable gases.

The second role that must be considered is that once char is formed, it can influence further degradation. When an adherent and insulating layer of char is built up on the surface of a polymer, this layer may insulate the underlying polymer from the flame and, thus, make further degradation more difficult. This assumes that the combustion of char is a difficult process and there will be some dependence on the composition of the char layer. The typical char layer consists of an amorphous carbon along the way to graphitization; the extent of graphitization will influence the combustibility of the char layer. Enhanced thermal stability may result if the char layer contains an inorganic component in addition to carbon.

CORRELATION BETWEEN CROSS-LINKING AND CHAR FORMATION

A cursory review of the literature would lead one to believe that any cross-linked polymer will be inherently less burnable than a non-cross-linked polymer (2). In fact, this is not the case. The resistance to combustion of a polymer is in some way connected to both the number of cross-links and to the strength of the bonds that make up the cross-linked structure (3). In recent work from these laboratories, it

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Char Formation

has been observed that cross-linked methyl methacrylates, copolymers prepared by the radical polymerization of various dimethacrylates with rective methacrylate, degrade at essentially the same temperature as does the homopolymer poly(methyl methacrylate). The most likely explanation is that the cross-linked structure is produced by very weak bonds that may be relatively easily cleaved thermally. Upon heating, the cross-linked atructure is lost and the resulting polymer is now no different from that of methyl methacrylate and its degradation proceeds in the same way (4).

III. FORMATION OF CHAR

It is believed that the temperature at the surface of a burning polymer is close to the temperature at which extensive thermal degradation occurs (usually 300-600°C). The bottom layer of char, near the polymer surface, is at the same temperature, whereas the upper surface, exposed to the flame, can be as hot as 1500°C (S). Therefore, fire-retardancy chemistry is concerned with chars, which may be produced at temperatures between 300 and 1500°C.

A polymer passes through several steps in the formation of char. These include (a) cross-linking, (b) aromatization, (c) fusion of aromatics, (d) turbostratic char formation, and (e) graphitization. Turbostratic char refers to an incomplete process of graphitization, when solid spheroids (precursors of graphite) appear in the molten carbonaceous material, typically at 500-700°C (6). At this point, the graphite layers are arranged in a parallel fashion, yet they are random in translation and rotation. The char formed during the combustion of polymeric materials is similar to the unbosuratic char.

in fire-retardant terminology, all polymers are usually classified as noncharable or charable, depending on whether or not they produce char under pyrolytic conditions (5). In terms of the chemical processes governing thermal degradation, polymers may be divided into three classes (7.8): (a) polymers that undergo chain scission and volatilize with, at most, a negligible amount of the char formation (e.g., polycthylene (PE), polypropylene (PP), polystyrene (PS), poly(methyl methacrylate) (PMMA)]; (b) polymers that undergo chain stripping reactions, producing unsaturation in the main chain with loss of hydrogen atoms and the pendant groups and give a moderate amount of char [e.g., poly(viny) chloride) (PVC), poly(vinyl alcohol) (PVA), polyacrylonitrile (PAN)]; (c) polymers that contain aromatic rings that can cross-link simultaneously with chain scission reactions and produce moderate to high amounts of char [e.g., aromatic polyamides (PA), polyesters (PS), polyearbonates (PC); polyimides (Pl)]. Different authoss may categorize each polymer in one or another category, depending on the emphasis they wish to place on a particular reaction.



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Leychik and Willde

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A. Polymers That Naturally Produce Char

Under certain experimental conditions, even aliphatic polyhydrocarbons can produce some char. The dissociation energy of C—C bonds in hydrocarbons is 65—90 kcal/mol, depending on the structure, whereas C—H bonds have a dissociation energy in the range 90—105 kcal/mol (9). Because these energies are similar, dehydrogenation may compete with the chain scission at high temperature (10), as shown Eq. (1):

Vinyl chain ends activate hydrogen in allylic positions; the dissociation energy of the allylic C—H is ~85 kcal/mol (5,9) and this leads to dehydrogenation and formation of conjugated dienes. The dienes may react with an activated double bond (dienophilic reagent) and lead to aromatization, as shown in Eq. (2):

The diene may also undergo further dehydrogenation leading to a triene that may cyclize [Eq. (3)]:

This is a phenomenological model of carbonization of polyhydrocarbons. However, under the normal degradation conditions in an inert atmosphere, the formation of char by these processes is quite limited. One of the goals of fire retardant science is to promote these char-forming mechanisms.

Polyacrylonitrite (PAN) has been studied very extensively (11) as a potential source for the production of carbon fibers. It was shown (11-13) that the thermal degradation process of PAN strongly depends on heating rate. Under rapid

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Changing Nature of Fire Retardancy in Polymers

ment of metals by plastics. It is not the cost of a material that determines material selection in an application, but what the final cost is to perform a function. As polymer options expand, as processing becomes more sophisticated, higher enginecred parts become more practical. One may thus find combustible organic polymers in functions where metals once predominated. That can lead to unanticipated hazards (5).

IV. WHAT DOES IT MEAN TO BE FIRE RETARDANT?

Fire retardancy or flame retardancy or "FR" can be very much in the eye of the beholder. Combustibility can depend as much on fire conditions as on polymer coroposition. Whether a material is an "FR" material is very much in the context of tests. Whereas designing laboratory tests to fully simulate real fire conditions is not possible, the increased use of lests which measure fundamental properties. then using those properties in mathematical models, is becoming increasingly successful and is the subject of discussions elsewhere in this volume. The behavior of materials in a fire can be described by several factors (6):

- 1. Pase of ignition-how readily a material ignites
- 2. Flame spread—how rapidly fire spreads across a surface
- 3. Fire endurance—how rapidly fire penetrates a wall or barrier
- 4. Rate of hear release—how much heat is released and how quickly
- 5. Ease of extinction—how rapidly or how easily the flame chemistry
- 6. Smoke evolution—amount, evolution rate, and composition of smoke
- released during stages of a fire Toxic gas evolution -- amount, evolution rate, and composition of gases released during stages of a fire

A multitude of fishmasbility tests measuring one or more of these properties have been developed. There are more than 100 tests in the United States alone. Other countries use different tests and rating systems, often making comparisons difficult. A material acceptable in one country may be rejected in another. A polymer resin may thus be more ignition resistant than its parent or than another reain and thus be "ignition resistant" or "flame returdant," but still have insufficient fire performance in the context of the tests appropriate for the application. That can lead to conflicts and miscommunication among resin supplier, customer, and end user. For example, in a recent legal case, a rubber manufacturer sold an ignition-resistant resin (UL-94 V-0) into an application. The application had its own industry test, which had few test facilities. The application test was known only vaguely to the resin supplier. The customer bought the resin, although it was

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marginal, because the resin was substantially less expensive than the alternatives. Years later in a product wide process change, the resin manufacturer found it necessary to reformulate its rubber. It formulated a new V-0 product which was not not elearly V-0 than the provious product. That "new improved product" even more clearly V-0 than the provious product. That "new improved product" was shipped to the customer under the same product number. Parts using the new resin were made and shipped. Only later did final product testing show that the new resin did not pass the end-product test. This necessitated a part recall from across the country, Litigation resulted. Who knew what? Who should have communicated what to whom? When? What quality control testing was needed and which would have shown that final parts did not comply with specifications? "Free retardant" is not an absolute. It has a context which must be understood by supplier, customer, and user alike. That context needs to be discussed and communicated clearly.

V. IS THE USE OF "FR" RESINS MEANINGFUL?

In 1988, the National Bureau of Standards reported on a fire hazard comparison of fire-retarded (FR) and non-fire-retarded plastics (7). Applications included a TV housing, a business machine housing, upholstered chairs, a cable array, and laminated circuit boards. In each case, a non-fire-retardant resin and a corresponding flame-retardant resin were used in the application. It was concluded that representative fire-retarded products showed significantly enhanced fire performance. They found the following:

1. The average available escape time was more than 15-fold greater for the FR products in room burn tests.

 The amount of material consumed in tests of the FR products was less than half the loss in the non-FR tests.

 Fire-retarded products, on average, gave one-quarter the heat release of non-FR products.

 The total quantity of toxic gas, expressed as CO equivalents, released by the FR products was one-third that for the non-FR products.

5. The production of smoke was not significantly different in room fire tests between FR and non-FR products.

"The study shows, then, that the proper selection of fire retardants can improve fire and life safety" (7). It was noted that such a conclusion should not be automatically expected, but that test and evaluation of candidate systems is necessary to ensure fire performance enhancement in the final application. It was also noted that production of carbon monoxide was not accurately predicted in less than room-sized tests. Insuruments like heat-release-rate calorimeters are overventilated versus some room test scenarios and thus substantially underpredict CO.

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P. 002

Chapter 8

Char Enhancing Approaches to Flame Retarding **Polymers**

J.W. Gilman, T. Kashiwagi, R.H. Harris, Jr., S. Lomakin, J.D. Lichtenhan2, A. Boifa and P. Jones2

Fire Science Division, Building and Fire Research Laboratory, National Institute of Standards and Technology.* Gaithersburg, MA 20899-0001 USA.

Air Force Research Laboratory, Edwards Air Force Base, CA 93524-7680 USA



ABSTRACT: Additives that increase the amount of charcoal-like residue or carbonaccous char that forms during polymer combustion are very effective fire retardants (FR) [1-3]. Our research efforts focus on reducing polymer flammability by promoting char formation. Our approach to char promotion is to investigate additives which enhance charring, and to gain a fundamental understanding of the additives' mechanism of char formation with the goal of optimizing their performance. Char formation reduces the amount of small, volatile polymer pyrolysis fragments, or fuel, available for burning in the gas phase, this in turn reduces the amount of heat released and fed back to the polymer surface. The char also insulates the underlying polymer, due to its low thermal conductivity, and reradiates incident energy away from the polymer surface. The char must also function as a mass transport barrier, by physically delaying the volatilization of decomposition products and/or chemically reacting with decomposition products. The physical structure of the char is important in this role. Thick, feating that appears to be more fire resistant than brittle, thin char [4]. This char-enhancing approach is most successful when the polymer chara rapidly and early in the burning process. To be useful, the charring process must occur at a temperature above the polymer processing temperature, but below the temperature where rapid gasification of the polymer to combustible fuel occurs. This chapter is an overview of several new char enhancing approaches to reducing the flammability of polymers.

8.1 Introduction

Current efforts in our laboratory focus on several different char enhancing, inorganic-additive systems. All of these systems entail the use of inorganic materials as additives to organic polymers, such as commodity polymers, engineering polymers, thermoplastics and thermosets. Under a class of additives best described as metal

135

Chemistry and Technology of Polymer Additives. Chapter 8. 1999; Blackwell Science Inc., Malden, MA, Ak-Malaika, S.; Golovoy, A.; Wilkie, C. A., Editors, 135-150 pp, 1999.

Exhibit D

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[†] Guest researcher at NIST from the Russian Academy of Sciences, Moscow, Russia.

Curtain commercial equipment, instruments, materials, services or companies are identified in this chapter in order to specify subquately the experimental procedure. This in no way implies endorsement or recommendation by NIST.

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NTRODUCTION

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be the basic literature source in this field of technology. Compilations of Journ articles in areas of spacific interest thus become a valuable literature resource, an to guide their content that within five years a complete collection of issues woul One of the goals of the Journal of Fire and Flammability and its supplements is i the Fire and Fiammability series of books embodies this concept.

Fabrics were the first materials in ow society to be the subject of recent fire safet concern, in large part because of the amounts and extensiveness with which it is present in everyday life. The Flammable Fabrics Act and its amendments expres the government's concern about the fire safety of fabrics.

erticles express this focus in their titles. Much of this information is novel am unique in coverage and viewpoint, and all of it is presented in the context of ou contain useful information on the flammability of fabrics, and fourteen of thes It is a measure of the importance of fabric flarunability, that twenty erlicks mission of helping to make this world safer for its people.

These twenty articles are presented in this volume, with the hope that it will indeed

prove to be a valuable literature resource in this field of knowledge.

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: WORLD

FIRE AND FLAMMABILITY SERIES VOLUME NINE

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(Received June 17, 1972)-

FLAMMABILITY AND FLAME RETARDANCY OF FABRICS: A REVIEW

ment of labric flammability, progress in the volence and rectinology of and glow retardation by the currently known liente and glow returdents. takics are critically reviewed with reference to the importance of rescercin ABSTRACT: Flammability and flams retardancy of the commonly used ilame returdation, theory of flams retardation and the mechanism of flar Several conduding remarks have been made. Thank are 68 reterances. on labric flame retardation; mechanism of flame prope

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INTRODUCTION

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involved in research and development to reduce or retard fabric Liammabillty, as is evidenced in the increased production and sale of flame relardants during the last craft, to name a few. The textile trade and altied industries are becoming more legistation have been summarized [1, 4, 5]. The concern of the consumer and the impact of regislation have shown the pressing need for flame retanded fabrics and thers for such applications as apparel, interior fumishings, automobiles, carpets, air Germany, Great Britain, Japan, Switzerland and the United Slates. Some of this tective measures can be seen in the development of legislation against the use of "Nighty or dangerously flammable fabrics" in countries such as Canada, France, as is evidenced by a meeting [4] having representatives trom many countries of the world. The consensus was that the consumer should be protected and such prosome role [4], the issue of tabric flammability has become an international concern [11-3]. It appears that, as estimated in the United Stales, 160,000-300,000 persons. the victims are young children. Because loss of life and property and severe suffering are often associated with burn injuries in which clothing materials play Relevant statistical wrveys have recently been reviewed per year suffer burn injuries as a result of clothing ignition. A high parcentage of Lb-low $300^{\circ}\mathrm{C}$, producing gaseous products; some of these are flammable and grite under lavorable conditions. Cansequently, fibers constitute fire hezards. Igniion of textiles causes perhaps about 20% of the total loss of life and property due, LMOST ALL TEXTILE fibers are organic polymers, most of which decompose opardactors) Remarch Assacietts. To whom correspondence should be addressed.

To Connie

P.05 #625

2003,05-21

FROM

Flammability and Plame Retardance of Fabrics

Table 1, Flemmability Characteristics of Fibers (trom Ref. 21)

Fiber	Tang.	Flame Temp. °C)	Flaremobility
Cotton	99	98	Burns readily with dian formation and alterplow.
Rayon	8	8820	Bure redily with ther formation and exterplow.
Acertate	476	380	Burns and meter sheed of flame.
Triacetate	2	885	Burns raddily and malts about of slame.
Nyton 6	93	\$75	Melts, supports combustion with difficulty.
Nylon 88	229	ı	Mella, does not readily support combustion.
Polyester	450	28	Burns resolity with melting and shoot.
Acrylic	98	5	Busns readily with meting and sputtering
Maderalic	i	1	Males, burns very slowly.
Polypropylena	570	S	Burns slowdy.
PVC and Polyvinyildens chloride Wool	† 69	1 \$	Does not support combustion. Supports combustion with difficulty.

gen and phosphorus content of the fabric. Miller et al. [27] studied the effects of various physical peremeters e.g fabric weight, specimen size, endosure chimney. horizontal and vertical mounting and direction of gas Illow on the OI and observed cardle like burning of the polymer. Jesoro et al. [10] found that Di measures values of some of the fabrics as measured by tester et al. tre shown in Table 2. Williard 126) found that the oxygen index is a simple function of the added altroflammability measurement of untreated and flame retarded fabrics. Originally developed by Ferrimore and Martin [26], the OI is defined as the minimum volume faction of oxygen in a slowly thing oxygen-nitrogen mixture that will sustain flammability of fabrics as a function of chemical composition of the liber. The Ol of the burning time to the charred length lor comparing the flammability of fabrics. extent. In racent years, the Oxygen Index (OI), method has gained popularity for but it his never been accepted and used for evaluation of flammability to any

M.A. Kasem and H.K. Rouette

M.A. Kasem and H.K. Rouette

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my a fabric speckmen of test of test of fabrics of fabrics. thods mentioned above have been critically discussed [12, 13]. It appears that fabrics may rank in different orders, even it the same-property is purportedly ng messured. Suchecki [24] suggested ån "Index of Merit" defined as the ratio ural and synthetic tabilics; the flame application time has also been reduced from sec to 3 sec [23]. Results obtained for flammability measurement by the test thod [16] has been widely used and has been recantly modified to test both lerms of attentiame (durotion for which a specimen flames after removal of the tion source), that length (vertical length of the specimen charred) and afterglow ration of glowing). Usually a test specimen is ignified by the application of a s prescribed conditions vary from method to method and also depend on the ulrements of the country concerned. The AATCC 34-1969 vertical char test diffed flame for a delimite length of time and the above criteria are measured. ess are shown in Table 1. The maximum Heme temperature of a fabric was found Test specimen (20) For flame retarded toprice, the flame reastance is evaluated e, "the maximum flame temperature" and the flammability [19-22] of common depend on specimen dimension and on the concentration of oxygen surrounding) the flame temperature (temperature inside the flame produced by a burning ric) are also measured for evaluation of fabric flammability. Ignition temperaimoplastic (abrics [18], Additional pharacteristics such as ignition temperature aric, the test specimen is given support either by the use of a backling or by ving of threads (usually glass fiber) along the specimen. Such arrangements have in found to give resonably good recults of flammability measurement on id as a criterion for flammability and such measurements on a series of fabrics w.been reported '9) The flammability of thermoplastic fabrics is evaluated by ire or less the same methods. However, in order to prevent melt dripping of the in addition, the rate of flame travel along an ignited fabric specimen may also be

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Figure 2. Schematic sepresentation of the burning

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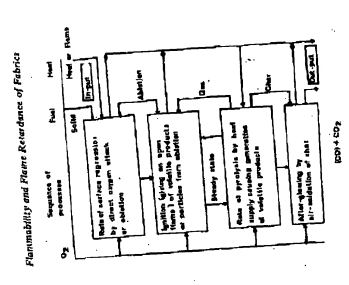
iew years [4]. The estimated market for flame retandant chemicals is expected to incresse from 200 million pounds in 1971 to 850 million pounds in 1975 [6] M.A. Keiem and H.K. Rouelle

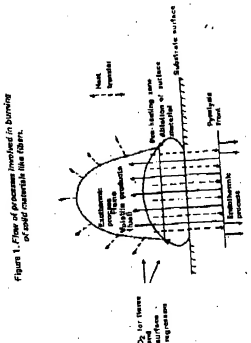
FLANMABILITY CHARACTERISTICS OF FABRICS

exygen index values for the blends with increase in the percent Namex in the Nomex with cotton and other flammable fibers, and they observed increased may reduce flammability of the flammable component of the blends, even to the seji extenguishing point. Tesoro and Rivlin [11] reported such a study on blends of blend from 0 to 15% and remains virtually independent of the cotton content (up to 85%) in the blend. Moreover, the presence of "nonflammable liberi" in blends (Ismmability of polyester increases with increase in the amount of polyester increases by the so-called "scaffold" affect [4]. Tesoro et al. [10] found that the artempted to relate fabric flammability with the above properties, and found that tegrier fabrics burn more slowly than the lighter onse. For blends, the flammability is influenced by the nature and proportion of the constituting fibers; in particular exothermic nature of the decomposition reaction becomes more pronounced with Increasing molecular weight. Fabric properties such as construction, weight per unit area and surface smoothness affect flammability. Nietson and Richards [9] weight of the fibrous polymer increases the decomposition temperature [8] and the cross-linking [7] decrease flammability because the fragments resulting during polymer decomposition are too large to be volatile. However, increasing malecular chamical linishing operations such as mordanting during chrome dyeing of wool i.e. The flammability of fabrics is a complex phenomenon and is influenced by the characteristics of both the finer and the fabric. The chemical composition of the fiber is the important litter characteristic affecting farmmability. In addition, Factors Affecting Fabric Flammability

Mechanism of Burning of Fabric

température. In contrast to surface regression, pyrolytis loyms a solid residue (char) thermic process, and results in the formation of volatile products. The latter diffuse out to the surface of the solid and form additional fuel, which increases the flame and the ablation. The heat which diffuses into the solid causes pyrolysis, an endofisme. Hence, the rate of surface regression depends on the direct oxygen attack surrounding stmösphere and/or diffuses into the solid. During the regression of the surface; absetion of solid particles into the flame occurs, thus acting as fuel for the surface of a Harramable material, the latter will ignite if a sufficient supply of exygen is evallable. The heet produced during burning causes convection of the but may be represented as shown in Figure 1. The process of ignition and burning of solid materials can be described schematically (Figure 2). If a flame attacks the The mechanism of burning 4combustion in air) is a complicated phenomenon





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Flammability and Flame Relardance of Fabrics

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M.A. Kotem and H.K. Rouelts

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daimed to be 1.4% [30].) Recently, Willer of al. [31] developed apparatus and gation rate! by moving fabric specimen at an appropriate velocity while maintaining the flame at a fixed position. It is claimed that data with a high degree of precision can be obtained by this method for both non-fusible and fusible fabrics, the fatter is more pronounced for flame retaided fabrics than for the untreated. The rapid and among laboratories. Reproducibility of measurement by OI technique has bren Lechnique "A wheel-led fame apparatus". For the determination of flame propa-Hendrix and coworkers [29] studied the effect of moisture content on the oxygen found that a moisture content of tess than 6% has very little effect. However, at a moisture content higher than 5%, the oxygen index values increase and this increase and intense interest in the DI technique is due to the pracision with which the OI values can be measured and the reproducibility of the residts within a faboratory index of colton, polyester and their blends, both untreated and liame retainded, and temperature of the text specimen on the OI of flame retarded cotton and found that the Ol decrease with increase in prejention semperature of the fabric. that they affect of significantly, DiPietra, [28] studied the effect of recignition without any support grid.

TECHNOLOGY OF FLAME RETARDANCY

Development of Flame Retardancy

Man's desire and need to reduce flammability of readity combustible materials go back to the prehistoric era, but attempts to impart flame resistance to fabrics are of relatively recent origin. In 1638 planer of Paris was used as flame retardant for or relatively recent origin.

canvas and in 1821 Gay-Lussac investigated a number of Inorganic salts as flam retardant for textile fabrics. From then until about 1950, a limited amount of a retardant for textile fabrics. From then until about 1950, a limited amount of a rate during the past 10-20 years. The early work was concerned primarity wit rate during the past 10-20 years. The early work was concerned primarity wit has summarized patent libratures on flame retardant textiles. The introduction of has summarized patent libratures on flame retardant textiles. The introduction of retardants for such fabrics. Some of the relevant literature have been reviewed literatures for such fabrics. Some of the relevant literature have been reviewed literatures used to impart theme resistance to polyester, mylon, acrylics and a fer retardants used to impart theme resistance to polyester, mylon, acrylics and a fer retardant theorem phosphorus, nitrogen and halogen. In recent years some work has been contain phosphorus, nitrogen and halogen. In recent years some work has been fabric [39, 40].

Ischniques of Fabric Flame Retardation

mongeners and siso-crosslinking sgents far celluloss. APO is capable of formit both home and copuly ner and such polymers have been formed on cotton t tardants are cheep, they are nondurable or semidurable to washing, depending c whether the retardant is soluble or insoluble in the washing solvent. Finishing (tabric with reactive type flame relandants offers fastness to stendard home washir and the effect is known as durable. Durable flame retardants for cotion fabr includes THPC-patymers, APO and APO-THPC, Both APO and THPC are potenti precipitated on cotton by the reaction of NaWO. and SnCl. The unreactive r (polymer) at some stage during application of the retardant to fabric. The us reactive class includes a host of inorganic and organic compounds, most of whic are saits. Water or organic solvent soluble compounds are applied from solution be most widely, used in textile field. Flame reverdants used for textile finishing me cation is based on whether or not the flame retardant reacts with the fabr be put into 180 segeral classes, unreachive and reactive. Apparently such a classif phosphonomethylation, phosphotomydation, mesylation, tosylation and cyani ethylation of caliusose and introduction of chlorine into polyester. In the copolmeizstion pracess one of the monomers contains a flame retarding element, e.t. chloiline, and is copolymetized with a second monomer so that the polymer cotains its flame retardant in the chain. Of the four approaches, linishing appears t while the insoluble compounds are formed in stru on the fabric, e.g., WOs ore: finishing fateirs with flame returding chemicals, adding flame retardants 1 chemicals and the product is flame retardant. Examples include phosphorylition polyment before their uttimate extrusion, copolymerization and chemical modific tion of liber. Chemical modification involves reaction of the polymer with som Several different approaches have been used fer flame relanding fabrics. Upon

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ponents of the blend. Since flame relandants are specific in their effect, it may be necessary to use more than one type of flame retardant for blends. Some of the recent studies in this area have been reported by Krisse [4]. somponent lakild, However, It is usually necessary to flame retain all the com-

Syneegism in Flame Retardancy

THPC-NH3, THPC-diamine etc. [43]. Such increase in the flame retarding efficiency is an example of synergism and is an attractive field of research both efficient frame retardants than the constituent compounds. Examples include Examples include brominated phosphates, TMPC-NM₃ polymer, and phosphonitrilic chloride. Similarly, certain mixtures of compounds have been found to be more boray boric scid, organic phosphate-ures, antimony exide-organic chloride, taining theme returdant has been found to increase the flame retarding efficiency. Imporporation of retrogen or halogen (F. Cl and Br) into a phosphorus comfrom theoretical as well as practical view points.

MECHANISM OF FLAME RETARDANCY

present knowledge is speculative. However, cellulose has been widely studied and a wealth of information is available. Several theories based on this information, some recent years to understand flame relardancy. It appears [49] that much of the eppicable to all polymer is yet to be evolved, atthough much effort is being given in The theoretical aspect of the mechanism of fabric flame retardancy is complicated both because of the complexity of the burning process (1self, which varies from polymer to polymer, and because of the diverse nature of the compounds used as flame retardants. A generalized and coherent theory of flams retardancy General

Flammability and Flame Retardance of Fabrics

of which is empirical and induitional, have been pur forward, but a systematic stud of the mechanism of flame retardancy in other polymers has begun only recentl [49]. However certain generalized datements pertaining to the nature of the flam

retainsants can be made.

mum" (10 percent) add-an is required to obtain a "satisfactory" flame of tardancy. This minimum add-on varies from polymer to polymer e.g. for 18 1. Plame retaining effect of a compound is "catalytic," elthough a "high min

same retardant viscose requires higher add-on than cotton.

factors such as thermal stability of both retardent and the polymer and chemic reaction at elevated temperature of the flame retardant with the basic polymi The effectiveness in terms of add-on of flame retardants depends on men

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Flams retardants are specific in their effect i.e. a compound may be a god flame retardant for cellulose but may not have any affect on polyester.

polymers e.g. cellulose, but their effectiveness does not appear to depend is niticantly on their structure. However, compounds decomposing above the Organophosphorus pompounds are especially affective for hydroxyl containing flaming temperature ($400^{\circ}\mathrm{C}$) are ineffective.

The efficiency of halogen containing flame setandants increases in the order 1 Cl. Br. I and halogens on aromatic structure are less effective than those on the ú

The Function of Sams Flame Retardants

aliphatics [50, 51]

flaming. Derivation of the function(s) of flame retardants is a formidable plece t work because of the many interacting variables involved in flame retardancy. Neve theless, some generalizations concerning the function of flame setædants pertainir In principle Hame relardancy can be achieved by affecting any or all of the stages of burning of polymers. These stages are (Figure 2) ignition, pyrolysis an primerity to celiutose (but perhaps applicable to other polymers) can be made.

1. Phosphorus containing flame retardants act predominately in the solid phase an affact pyrolysis of the polymer.

affect the flaming reaction by acting as free radical scavengers; they may siso at 2. Halogen containing flame retardants act predominately in the gas phase an

in the solid phase or in both phases.

...j. Differential thermal analysis (OTA) shows that (tame retardants lower the ch composition temperature of certain polymers [28, 39, 52.55]

3. Thermogravimetric analysis (TGA) shows that many flame retardants affect it pyrolysis in such I way as to produce more that and less flammable gas; maje weight loss of flame retardant fabric occurs at higher temperature compared t the untreated fabric (28, 53-55)

P.09

2003.05-21

FROM

Figurnability and Flame Resardance of Pabrics

M.A. Kasem and H.K. Rouette

"The non-combustible gates produced may reduce the exothermic surface re-5. Flame retardants may lower the heat of combustion of the polymer because less combustible gasss are produced; this in turn affects pyrolysis of the palymen. gression because they dilute the oxygen necessary to attack the surface

Finne retardants may form a coating film around the fiber such that oxygen on decomposition, produce non-flammable gases which dilute the pyrobysis necessary for flaming connot attack the surface directly. Flame reterdants may, products to a concentration below that which is required to sustain flaming. æ

Flame retaidants may act as inhibitors for the free radical (laming reaction e.g. halogens, acting as free radical scavengers in the flaming reaction.

Pyrolysis and Flame Retardancy

that the phosphorus compounds are converted to phosphoric or polyphosphoric the case of celtulose and phosphorus containing flame retardants, it is speculated nature of both the polymer and the flame retaidant and their thermal stability. In flame becomes unstable and may extinguish, lift: off or flash back outside the Hammabilly limits. The manner in which flame retardents affect the pyrobysis of polymers and the nature of interaction depends, among others, on the chemical fuel to oxidizer ratio, called the lower and upper fearmability limits [59]. The retarded cotton, although the proportion of the fishmable gases in the latter is . A combustible mixture will undergo flaming reaction only over a narrow range of reduced. The effect of this increased or decreased availability of pyrolysis products (fuel) is to preate an imbalance in the fuel to oxidizer ratio in the flaming reaction. retardent cellutose, and probably most other tabrics, produces more char and less [67, 58]. A recent mass spectrometric study [58] shows that the pyrohysis products from unitested cotton do not differ qualitatively from those of flame discussed here. It appears that all work leads to a common conclusion that flame flammable gas. In addition, flame retardents catalyze the pyrolysis of a polymer published [56] covering the basic aspects of cellulose pyrolysis which will not be interest to study the effect of flams retardants on pyrolysis. With the exception of a few, the reported work is related to cellulose and an excellent review has been . Pyrolysis of the polymer supplies fue: for flaming; hence it has been of utmost acid which acts as dehydration catalyst for cellulose [49].

Synergism in Flame Retardancy

thermal reaction of the constituent compounds. For the P-N synergism, Hendrix [60, 61] has shown that some organic phosphate in the presence of unea forms phosphoramide and the latter acts as flame retardant for cellulose. The studied. It is felt that the mechanism of synergism is depicted primarily by the little progress has been achieved. Only a limited number of systems have been Basic aspect of synergism is a relatively new tield of investigation and hence very

halide, which acts as tiams retardant [50] . Similarly phosphotus-halogen syne detailed disoussion of such synergistic effects can be found in the recent issue antimony-halogen synargism is thought to be due to formation of antimor is thought to be due to formation of phosphorus -tri or -penta halide [5] of the Polymer Conference Series.

Dehydration Theory of Flams Retardancy

are applicable only to a limited number of flame retardants. The chemical ti retardants. Such a theory has yet to be evolved. However, several theories have put forward for cellulose. These ere: the coating, the gas, the thermal an In principle a theory should be able to explain the mechanism of acti chemical theories and they have been described elsewhere [62]. The former known fiams retardants and in addition, should be able to predict new appears to be more general and applicable to a large number of compounds.

returdants are either Lewis acids or bases or their precursors and are available near the flaming temperature of cellulose to catalyze dehydration of cell through a carbonium of a carbanion as shown below. Such a dehydration pr may continue until all the "available" hydroxyl groups of cellulose have The chemical theory was developed by Schuyten [63]; it assumes that involved, leaving a residue of high carbon content (char).

FR . Fleme Rebrutfi H-C-OH + FMACIO) -----N-C-A-C-A-C-FH) - 13-0 + FR. __ (#2-HO)+eH

GLOW RETARDANCY

HAO +FR

15:22

oxidation of carbon to cerbon monoxide is less exothermic than that to ca carbon diaxide. Thermolebile phosphorus and boton containing compounds some metallic oxides have been found wetul as glow retardants [64, 65]. Se postulations have been put forward to explain glow retardancy of cettolose. dioxide and it has been proposed [66] that glow retardants divert oxidation Glowing (flameless combustion) is due to the oxidation of carbon (cha

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CONCLUDING REMARKS

phosphorus containing glow intardants were proposed to act as inhibitors for the [64-66] so that oxygen can not diffuse into the char. Furthermore, boron and

carbon to the less exathermic carbon monoxide so that the heat of exidation is posed that glow reterdants may coat the "active exidation sites" in the char insufficient to sustain further oxidation of casbon. In addition, it has been proexidation of carbon by interfering with the electron transfer process in the exida-

made to approach this goal. Navastheless the current test methods provide data which may be used to classify fabrics as slow or fast burning, flammable, less fabrics in actual use is far from a reality, although efforts are cominworshy being sammability of fabrics, but a method which can simulate the flammability of The need and importance for flame retarding fabrics are increasing at a faster rate than ever before. A large number of test mathods are available for avaluating

flams retardation of cellulose. Studies on synthetic libers and their blends with natural fibers have been neglected in the past. In view of shelr increasing use, more siforts should be directed towards understanding basic aspects of flammability and flame retardation of synkhetics and blends, and it appears that activity in this area is tardation of fabrics other than callulose which shares most of the literature in this field and much is known about the basic and applied aspects of flammability and Very little is known about the basic espects of Hammability and flame reflammable or more flammable under specified conditions.

is an interesting, chaltenging and fruitful field for Investigation and it remains up to Fiammability and tlame retardation of polymenic materials, particularly fabrics, properties and the sesthetic values of fabrics. Only from the results of systematic studies, may it perhaps be possible to tailor flame retardants which would meet home laundering. In edottion, most flame retardard finishes affect both the physical Although a large number of flame retardants are known particularly for cellulosic fibers, ordy a very few of them posses any reasonable fastness to normal most if not all the requirements asked for by the consumers. gaining momentum.

scientists to explore and develop this rather unsystematic and unorganized branch of science and technology.

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tion of carbon [85]

FROM

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(Ascelved January 14, 1970)

namnability in oxygen-eartched atmosphere been conducted. Burning rate data for a standard material in Studies of the effects of environmental parameters upo He resistance for measuring flammability In hyperbanc environments and nimgen and oxygan-hallum mixtures are

INTRODUCTION

absolute oxygen concentration and the oxygen partial pressu quantity of nammable materials in the chamber should be red tion. In practice, it is often difficult or impossible to remove an probability of further applicants in oxygen-enriched environme probability of further any one of the three fundamental prefequis arrive to eliminate any one of the three fundamental an ignition source, a fuel and an atmosphere capable of suppo requirements with absolute certainly, Attempts should therefor reduce the contributing effects of all three besemine factors. associated with the use of these environments. Although fires bers has politied up to the vital need for research into the combi and hypoteric chambers are rare, they have occasionally occur pital hyperbaric chambers, oxygen tents and divers' decomp shing suddentess and often tetal results [8, 7 8, 9]. In order ever-increasing application of oxygen-enriched atmospiliems that may serve as polential ignition sources should chamber should be minimized as much as possible.

material, (2) determine the minimum exygen concentration re-(3) catagorize the tlammability hazards of cloth fabrics, elastom effects of environmental parameters upon the combustion rate combustion of solid flammables in the presence of nitrogen ar in order to evaluate the influence of the above factors upon oxygen-enriched environments, a study has been conducted to tions in hyperbanc environments.

Fire Studies in Oxygen-Enriched Atma

. An osygan-emichad atmosphere (DEA) is generally delined as an amosphere will than 0. than 21 mole-% oxygan usofos contains a purital presente of oxygan usofos contains a

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FROM :

Flammability Measurements and Thermal Decomposition of Textiles

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(Received March 16, 1919)

Caroline Narrow Fabrics. Raybestos-Manhaltan

Dynatech Dies Posit

> (Fiberglass/asbastos composite) Kepton electric wire insulation

Carborundum

INTRODUCTION

has led to the generalization that "most or all flame-relardant linishes rep-A survey of the technical literature related to tlame-retardant compo When considering flammability, it must be remembered that virtually all terials are flammable in certain atmospheres and conditions. Even m and ceramics flame when exposed to extreme conditions. Therefore, mability becomes a relative, rather than an absolute, matter. There is no thing as a fireproof fabric, but there are different degrees of fire resistance.

· Several types must be heat treated (800°F for several hours) to remove organic resi-

dues before qualitying for Class 9.

Insulation Electrical

HISTORICAL

into nonflammable vapors on healing (2). Since Gay Lussac's investig of extensive literature pertaining to flame relardants has become available of extensive inerature penaliting to tights of more practical than theor a fortunately, much of this information is of more practical than theor a fortunately, much of this infittle application to a description of lunding value, and consequently it is of title application. phosphate by 1786. Gay Lussec completed the first systematic investigation he concluded the use of flame retardants in 1820. From this investigation he concluded the use of flame retardants in 1820. tal mechanisms of pyrolytic degradation and itame relardation. In the years, with the use of chromatographic and thermal analytical techniqu the study of thermal decomposition, fundamental data have been collected alum was found to give textiles tlame-retardant properties, as was ammo of covering the material with a glassy layer or those which would decom Application of tlame relardants to textile materials had its beginning t the most effective salls were those which were either low melting and ca 1640 when theater canvas was treated with clay and plaster of Parls. By

to be effective consist of insolubilized organophosphorous polymers coring nitrogen" [1]. Such generalizations have fed to common acceptant the theory that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts a synergistic effect on the flame-relative that nitrogen exerts are not not nitrogen exerts. properties of phosphorous-containing compounds.

Owens-Coming Supplied by B. Welson Supplied by B. Welson

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Sent By: FKGGB*FTL--KAIN;

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E. Hundrk, T. K. Anderwn, T. J. Clayton, E. S. Okon, and R. H. Barker

Sent more quickly than the undreated fabric. While the total amount of

sitile products may be tower with A then with B, they are produced

sitile products may be tower with A then with B, they are produced

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sitile products may be tower with A then with B, they are produced

sitile products may be tower with A then with B, they are produced

to the products may be tower with A then with B, they are produced

at the A because the finish catalyzes the decomposition of the same and name temperature and nonburnable the same the same temperature state of a to be very careful when defining than retardance. Some obvious is to be very careful when defining than cataly the result means that a to be very careful when defining than cataly the same blands. Each of the same of accessibility of feat to the liber surface or presence of a compensature of the fabric. This may determine a tabric's ease of burning catarogehere.

Leathogehere

subjected, will influence its flammability.

to liaming. The results of any single test would be difficult to correlate is test indicates that no single test will effectively define a tabric's resiste subjected, will influence its fiammability.

nd use.

uld also be noted that the evidence for the presence of pellulose is changed during pyrolysis because the entire mode of decomposition A, The lowest point of the 160% cotton and otherm is at 360°C, while west point of the dyed cotton endotherm has been shifted to 320°C beed the effects of sulfur dye. However, the analysis indicates that these is of the tlame retardant finished sample indicates the presence of two odes of decomposition remain essentially the same. On the other hand the results obtained when three different botton samples were studied sidues during the combustlen of a given earnple. Every cotton earnple tod shows that the decomposition temperature of cotton is lowered en adulterant is present in the labric. The adulterant may be a softener, ill, flame resistant linish or residues not removed during the preparation ss. Differential Thermal Analysis (DTA) studies show that not only is the ise is changed when resistance to taming is achieved. Figure 9 Illussature differential importent, but that the mode of decomposition of ernal analysis techniques can determine decomposition temperatures strins, one milnor and one major, at 313°C and 267°C, TOA And Oxygen Demend

nable and provides the residue data which is of importance. Figure 10 ere analyzed, and Table 3 provides the consolidated data for these same pse results show that the addition of products lowers the onset decomermogravimetric Analyzer (TGA) data support the change of temperature the progression of residues and temperatures when five different fab-

in temperature and that the percentage residue varies with residual me-

ئے۔ و and a 20-C/ Bere the skipe is +0.2 dyed cotton fabrics with bisached cotton in an alt absorphere comparing THPOH treated collon and audio Vecour 201 using a 2 num capillary tube, glass reference, ş Per ich <u>भ्रेल</u>श् 300 minute heating rate starting of 25°C. Gecou More and at is 0,5-1 division 12446 -1

cotton fabric and THPOH Figure 10. TGA of puta cellulose, propared cotton labric, sulusing 1/2. Hariminate altrogen atmosphere, 0 shift, 5-5/minute heating rate and 4 mg/division ver T't ifteen and 45 tur dyed cotton fabric, yet treated var dyed tabric

19.67 19.67 21.17 ģ Table 3. Onset Decomposition Temperatures, Petcent Residues and Oxygen Demand 0.83 9.18 12.90 13.46 **1**2 o T 386 315 375 275 Data for Salacted Cotton Febrics Unitreated Sulfur Dya Control Univested Vat Dya Control. repaired Cotton Fabric THPOK Vet Dye (30%) Pure Cartion Celluloss

A major residue of the suffur dyed tabric is associated with B tem-ire deflection of 65 degrees below that of pure cotton cellulose, even